

Python Practice and Workshop 1

Printing data

Practice printing out data

Reading from a file and counting information from a file

Let's count up Exon number and length.

Use `/bigdata/gen220/shared/simple/rice_random_exons.bed`

Write python code to do the following:

1. Open the file
2. Loop through each line in the file
3. Print the length of each exon
4. Summarize the total length of all the exons

Reading from a comma delimited file Use the data file from this

site: <https://datacarpentry.org/2015-03-09-ISI-CODATA/data/biology/species.csv> for this example.

Print out the names of all the genera (genus column) and the counts of each.

FASTA file processing

Let's read in a dataset of Genes from a FASTA formatted file and print out: 1. The number of sequences in the file. 2. The first and last codons (first 3 bases and last 3 bases) 3. Count the number of genes on each strand (this is coded by the last character in the gene name).

```
import itertools, sys, re

# define what a header looks like in FASTA format
def isheader(line):
    return line[0] == '>'

def aspairs(f):
    seq_id = ''
    sequence = ''
    for header,group in itertools.groupby(f, isheader):
```

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        if header:
            line = next(group)
            seq_id = line[1:].split()[0]
        else:
            sequence = ''.join(line.strip() for line in group)
            yield seq_id, sequence

filename="/bigdata/gen220/shared/data_files/S_cerevisiae_ORFs.fasta"
#filename="S_cerevisiae_ORFs.fasta"
with open(filename,"r") as f:
    seqs = dict(aspairs(f))
    first_codon = {}
    last_codon = {}
    sequence_count = 0
    strand_count = {}

    for seqname in seqs:
        sequence_count += 1

        firstcodon = seqs[seqname][0:3]
        lastcodon = seqs[seqname][-3:]
        if firstcodon in first_codon:
            first_codon[firstcodon] +=1
        else:
            first_codon[firstcodon] = 1

        if lastcodon in last_codon:
            last_codon[lastcodon] +=1
        else:
            last_codon[lastcodon] = 1

    #     print(firstcodon,lastcodon)
    last_char = seqname[-1]
    #     print(last_char, " in ", seqname)
    if last_char in strand_count:
        strand_count[last_char] += 1
    else:
        strand_count[last_char] = 1

print("1.")
print("There are %d sequences in the file"%(sequence_count))

print("2.")
print("The distribution of first codons is:")

for codon in first_codon:
    print("%s => %d (%.1f%%)" % (codon, first_codon[codon],

```

```

100.0 * first_codon[codon] / sequence_count))

print("The distribution of last codons is:")
for codon in last_codon:
    print("%s => %d (%.1f%%)" % (codon, last_codon[codon],
100.0 * last_codon[codon] / sequence_count))

print("There are %s genes on the Watson (+) strand"%(strand_count['W']))
print("      %s genes on the Crick (-) strand"%(strand_count['C']))

print("3.")
print("There are %s genes on the Watson (+) strand"%(strand_count['W']))
print("      %s genes on the Crick (-) strand"%(strand_count['C']))

```